PATENT APPLICATION

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicants : James W. Baumgartner et al.

Serial No. : 09/090,867

Filed : June 4, 1998

For : TESTIS-SPECIFIC RECEPTOR

Examiner : Lazar-Wesley, E.

Art Unit : 1646

Docket No.: 95-33D1

Date : July 21, 1999

Assistant Commissioner for Patents Washington, D.C. 20231

Declaration Under 37 C.F.R. § 1.131

Sir:

We, James W. Baumgartner, Theresa M. Farrah, Donald C. Foster, Frank J. Grant, and Patrick J. O'Hara, do hereby declare as follows:

- 1. We are the inventors of the above-identified patent application.
- 2. All of the work described herein was performed in the United States of America by us or under our direction.
- 3. We have reviewed laboratory notes and other records, including the exhibits submitted herewith, and have determined that the invention recited in claims 1-32 of the above-identified patent application was reduced to practice before March 1, 1996 or was conceived before March 1, 1996 and was subsequently constructively reduced to practice with the filing of the patent application on March 13, 1996.
- 4. Attached hereto as Exhibit 1 is a copy of a computer printout of the DNA and deduced amino acid sequence of a clone designated "zcytor2." This printout is dated

prior to March 1, 1996. The sequences shown in Exhibit 1 correspond to those disclosed in the patent application in SEQ ID NO:1 and SEQ ID NO:2.

- 5. Attached hereto as Exhibit 2 is a copy of a portion of a memo written by one of us (Frank J. Grant) before March 1, 1996, which describes particular goals for the WSXWS receptor project, which project included the zcytor2 receptor. As stated in the memo, these goals included preparation of soluble forms (i.e., extracellular ligand-binding domains) of receptors. The memo also describes our intent to clone and express full-length, receptor-encoding cDNAs.
- 6. Attached hereto as Exhibit 3 is a copy of a page from the notebook of Cameron Brandt, a research associate working under our direction. This page, written before March 1, 1996, describes a plan to prepare polypeptide fusions comprising a soluble receptor and an immunoglubulin Fc polypeptide.
- 7. Attached hereto as Exhibit 4 is a copy of a slide prepared by one of us (Donald C. Foster) for an inhouse seminar on the WSXWS receptor project. This slide was prepared before March 1, 1996. This slide illustrates a plan to express new receptor-encoding DNAs in cultured cells, whereby the cells would produce the encoded receptor.
- 8. On the basis of these Exhibits we conclude that the invention recited in claims 1-32 of the patent application was reduced to practice before March 1, 1996 or was conceived before March 1, 1996 and was subsequently constructively reduced to practice with the filing of the patent application on March 13, 1996.

We further declare that all statements made herein of our own knowledge are true and that all statements made on information and belief are believed to be true; and further, that these statements were made with the knowledge that the making of willfully false statements and the like is punishable by fine or imprisonment, or both, under

Section 1001 of Title 18 of the United States Code, and may jeopardize the validity of any patent issuing from this patent application.

James W. Baumgartner	Date
Theresa M. Farrah	
Donald C. Foster	
Frank J. Grant	
Patrick J. O'Hara	Date

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HZCYTORO2.SEQ -
 Sequence of pcr products generated with 9800-9802,
 nested pcr product 9941-AP2 (9801-AP1)
 nested pcr product 9937-AP2 (9803-AP1)
 Enzyme Recognition
                    Cut Site
 Agel (A^CCGGT)
                     Def: 1124
. BamHI (G^GATCC)
                     Def: 172
 Drai
       (TTT^AAA)
                     Def:
                          36
 EcoRI (G^AATTC)
                     Def: 450
 EcoRV (GAT^ATC)
                     Def: 438
 HpaI
       (GTT^AAC)
                     Def: 145
 MscI
       (TGG^CCA)
                     Def: 1244
 MunI
       (C^AATTG)
                     Def: 493
 Ncol
       (C^CATGG)
                     Def: 377
 Nsil
       (ATGCA^T)
                     Def:
                          592
 Ppu101 (A^TGCAT)
                     Def: 588
      (CCC^GGG)
 SmaI
                     Def:
                          11
       (AAT^ATT)
 SspI
                     Def: 503 988 1107
Xma I
       (C^CCGGG)
                     Def:
HZCYTORO2.SEQ Linear
                             LENGTH = 1289
            Xmai
                                    Drai
   1 CCCCCCGCCCGGGAGAGAGGCAATATCAAGGTTTTÄAATCTCGGAGAAATGGCTTTCGTTTGCTTGGCT
     GGGGGGCCCCCCCCCTTATAGTTCCAAAATTTAGAGCCTCTTTACCGAAAGCAAACGAACCGA
                                               MAFVCLA
  70 ATCGGATGCTTATATACCTTTCTGATAAGCACAACATTTGGCTGTACTTCATCTTCAGACACCGAGATA 138
     TAGCCTACGAATATATGGAAAGACTATTCGTGTTGTAAACCGACATGAAGTAGAAGTCTGTGGCTCTAT
     IGCLYTFLISTTFGCTSSSDTEI
          KpaI
                                  BamHI
 139 AAAGTTÄACCCTCCTCAGGATTTTGAGATAGTGGATCCCGGATACTTAGGTTATCTCTATTTGCAATGG 207
     TTTCAATTGGGAGGAGTCCTAAAACTCTATCACCTAGGGCCTATGAATCCAATAGAGATAAACGTTACC
     K V N P P Q D F E I V D P G Y L G Y L Y L Q W
                                  172
 208 CAACCCCCACTGTCTCTGGATCATTTTAAGGAATGCACAGTGGAATATGAACTAAAATACCGAAACATT 276
    GTTGGGGGTGACAGAGACCTAGTAAAATTCCTTACGTGTCACCTTATACTTGATTTTATGGCTTTGTAA
    Q P P L S L D H F K E C T V E Y E L K Y R N I
 277 GGTAGTGAAACATGGAAGACCATCATTACTAAGAATCTACATTACAAAGATGGGTTTGATCTTAACAAG 345
    CCATCACTTTGTACCTTCTGGTAGTAATGATTCTTAGATGTAATGTTTCTACCCAAACTAGAATTGTTC
    GSETWKTIITKNLHYKDGFDLNK
                                Ncol
 346 GGCATTGAAGCGAAGATACACACGCTTTTACCATGGCAATGCACAAATGGATCAGAAGTTCAAAGTTCC 414
    CCGTAACTTCGCTTCTATGTGTGCGAAAATGGTACCGTTACGTGTTTACCTAGTCTTCAAGTTTCAAGG
    GIEAKIHTLLPWQCTNGSEVQSS
                         EcoRV
                                   EcoRI
 415 TGGGCAGAAACTACTTATTGGATÁTCACCACAAGGÁATTCCAGAAACTAAAGTTCAGGATATGGATTGC 483
    ACCCGTCTTTGATGAATAACCTATAGTGGTGTTCCTTAAGGTCTTTGATTTCAAGTCCTATACCTAACG
    WAETTYWISPQGIPETKVQDMDC
                         438
                                   450
            Muni
                     SspI
 484 GTATATTACAATTGGCAATATTTACTCTGTTCTTGGAAACCTGGCATAGGTGTACTTCTTGATACCAAT 552
    CATATAATGTTAACCGTTATAAATGAGACAAGAACCTTTGGACCGTATCCACATGAAGAACTATGGTTA
    V Y Y N W Q Y L L C S W K P G I G V L L D T N
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493

503

DI
NSTI

1553 TACAACTTGTTTTACTGGTATGAGGGCTTGGATCATGCATTACAGTGTGTTGATTACATCAAGGCTGAT 621
ATGTTGAACAAAATGACCATACTCCCGAACCTAGTACGTAATGTCACACAACTAATGTAGTTCCGACTA
Y N L F Y W Y E G L D H A L Q C V D Y I K A D
592
588

622 GGACAAAATATAGGATGCAGATTTCCCTATTIGGAGGCATCAGACTATAAAGATTTCTATATTTGTGTT 690 CCTGTTTTATATCCTACGTCTAAAGGGATAAACCTCCGTAGTCTGATATTTCTAAAGATATAAACACAA G Q N I G C R F P Y L E A S D Y K D F Y I C V

691 AATGGATCATCAGAGAACAAGCCTATCAGATCCAGTTATTTCACTTTTCAGCTTCAAAATATAGTTAAA 759
TTACCTAGTAGTCTCTTGTTCGGATAGTCTAGGTCAATAAAGTGAAAAGTCGAAGTTTATATCAATTT
N G S S E N K P I R S S Y F T F Q L Q N I V K

760 CCTTTGCCGCCAGTCTATCTTACTTTTACTCGGGAGAGTTCATGTGAAATTAAGCTGAAATGGAGCATA 828 GGAAACGGCGGTCAGATAGAAATGAGCCCTCTCAAGTACACTTTAATTCGACTTTACCTCGTAT P L P P V Y L T F T R E S S C E I K L K W S I

829 CCTTTGGGACCTATTCCAGCAAGGTGTTTTGATTATGAAATTGAGATCAGAGAAGATGATACTACCTTG 897
GGAAACCCTGGATAAGGTCGTTCCACAAAACTAATACTTTAACTCTAGTCTCTTCTACTATGATGGAAC
P L G P I P A R C F D Y E I E I R E D D T T L

898 GTGACTGCTACAGTTGAAAATGAAACATACACCTTGAAAACAACAAATGAAACCCGACAATTATGCTTT 966
CACTGACGATGTCAACTTTTACTTTGTATGTGGAACTTTTGTTGTTTACTTTGGGCTGTTAATACGAAA
V T A T V E N E T Y T L K T T N E T R Q L C F

SspI

1036 TGCTGGGAAGGTGAAGACCTATCGAAGAAAACTTTGCTACGTTTCTGGCTACCATTTGGTTTCATCTTA 1104
ACGACCCTTCCACTTCTGGATAGCTTCTTTTGAAACGATGCAAAGACCGATGGTAAACCAAAGTAGAAT
C W E G E D L S K K T L L R F W L P F G F I L

SspI AgeI

1105 ATATTAGTTATATTTGTAACCGGTCTGCTTTTGCGTAAGCCAAACCCTACCCAAAAATGATTCCAGAA 1173
TATAATCAATATAAACATTGGCCAGACGAAAACGCATJCGGTTTGTGGATGGGTTTTTACTAAGGTCTT
I L V I F V T G L L R K P N T Y P K M I P E
1107 1124

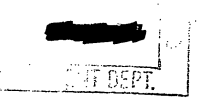
1174 TITTTCTGTGATACATGAAGACTTTCCATATCAAGAGACATGGTATTGACTCAACAGTTTCCAGTCATG 1242
AAAAAGACACTATGTACTTCTGAAAGGTATAGTTCTCTGTACCATAACTGAGTTGTCAAAGGTCAGTAC
F F C D T .

MscI

1243 GCCAAATGTTCAATATGAGTCTCAATAAACTGAATTTTTCTTGCGAA 1289 CGGTTTACAAGTTATACTCAGAGTTATTTGACTTAAAAAGAACGCTT

1244

DRAFT



Outline of things to consider for patent application of novel type I cytokine receptors

We have identified partial cDNA sequences for three new members of the type I cytokine receptor family. These receptors are characterized by a conserved cysteine pattern and an amino acid motif containing WSXWS. Members of this family include the receptors for TPO, EPO, Growth Hormone, Prolactin, IL-4, IL-7, IL-9, IL-2, IL-5, IL-3, GM-CSF, IL-6, CNTF, G-CSF and Leukemia inhibitory factor.

The main utility for these sequences would be to facilitate the cloning of the unknown ligands for the receptors. The receptors themselves (ie. soluble forms) might be potential therapeutics as well.

There are at least three ways the receptor sequence can be utilized to clone the ligands:

- a). Make receptor dependent cell lines (as was done in the project) for use in an expression cloning project.
- b). Soluble forms of the receptor can be labeled and used as probes in an expression cloning system.
- c). The receptor could be attached to various columns or other supports and used to purify the ligand.

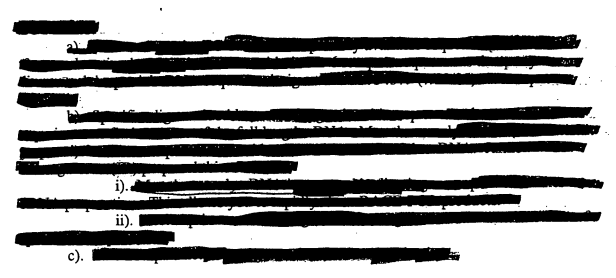
Patentable entities: (???????)

- a). The EST (expressed sequence tag) that allowed us to identify the partial ce as novel member of the family.

 i). Allows us to clone the full length and the full le sequence as novel member of the family.

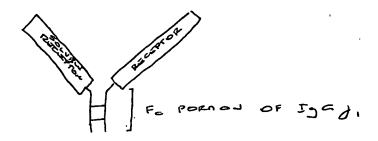
 - b). The full length receptor encoding cDNA.
- (c)/ Homologues of the cDNAs. It may be that murine versions of these receptors are necessary for ligand dependent cell line cloning.
 - d). The ligands for the receptors.
 - e). AIDS therapies. Discuss w/ Frank

WHAT WE GOT:



PURPOSE: WILL BUILD A VISCOR FOR EZPRITS: OF SOURCE
PRECIPTORS: FUSISD TO ISC J. HOWIT CHA. J. MIS
EXPRISSION SISTEM ALLOWS AN EDEN NAM TO PURPOS
SOURCE RECENTOR OWER A PROTECULA A COMMA

THOU PRESIDES A HANDLE POR VINC IN COING
AFTER LICANO.



- IJG PORMON OF RESON INCLUDES HAVE PLESONS CH2 ! CH3
- RUSSON IS CHERCUSO AS A MONOMOR BUT DIMERIZES

VIA IT'S TWO CHERCUS'S IN THE HAVAI REGION

لسفياح

5':

Ara SER.

GLU-PRO-LAS-SER-CTE-ACP-LAS-THE-HS-THE...

CONSTRUCTION OF BOILT SINE (FAUSCON STAL.

TO ELIMINATE UNBOUND

CIS IN HOUR PERIOD.

THIS CIS NORMAN BIND

LIGHT CHAIN, BUT LIGHT

CHAIN IT NOT NECESSARY

FOR CREATION OF A

FUNCTIONAL PUSION

(1300NETT, ET A. J. OF BIOL.

CHAINSTEPP 266 (34) 23060-23067

DET E, 1991)

3': WILL BE IDENTICAL TO NAM JE

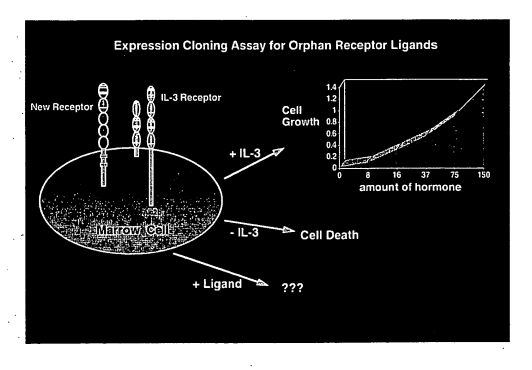


EXHIBIT 4